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Introduction

Nanotechnology is an exciting area of research involving nanomaterials with sizes in the 1–900 nm range. It has an everincreasing number of applications that cover a wide range of industries, such as medicine and life sciences. A promising application is its utility in biosensors. For instance, medical researchers have leveraged the benefits of nanotechnology in areas ranging from early detection of Alzheimer's disease to diabetes management to early cancer screening, including Covid-19 drug discovery.^{1–3} Nano-scale sensors are utilized in cases requiring greater precision and continuity.^{4–7} Moreover, they play an important role in the development of the emerging initiative bio-digital twins.^{8,9} Cancer is the second leading

Ultrasensitive low-probe-concentration PANC-1 and MCF-7 cancer cell sensors enabled by combined 2D-material-polymer-phage frameworks[†]

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Unique biophysical properties of cancer cells can provide essential knowledge that can aid in the diagnosis of disease, clinical trial designs and drug developments. The collective response of cancer cells is of significant interest to the research community. However, sensing cells in a medium population is extremely challenging. Here, we controlled the electrical conduction of cancer cells by using combined 2D-material-polymer-phage frameworks *via* current flow effects. We then developed an AC-pulse sensor system based on MoS₂/PEG/M13 nanoprobes for accurate detection of clinically relevant cancer cells such as PANC-1 cells and MCF-7 cells. The detection limit can reach ~15 cells per μ L, which is below an average of ~45 cells per μ L for existing sensing methods with medium cell populations. Moreover, excellent cell viability was achieved in PANC-1 cells for nanoprobe concentrations of up to 50 vol%, which surpassed those used in detection experiments. Furthermore, a nanoprobe concentration of ~10 vol%/7.1 nM was achieved, below an average of ~195 nM used for state-of-the-art sensing methods with medium cell populations. The combination of a previously unreported exotic sensing material and an AC-pulse strategy represents an approach for unlocking the ultrasensitive detection of cancer cells and provides a promising avenue for early cancer diagnosis, staging and monitoring.

cause of death worldwide, accounting for 10 million deaths in 2020.¹⁰ The cancer burden continues to grow globally. Nevertheless, survival rates for many types of cancers are improving, thanks to accessible early detection.

Recently, characterizing electrical signatures of cancer cell populations has generated substantial interest in the scientific community.^{11–16} The unique biophysical properties of each cancer cell type provide essential knowledge that could aid in the diagnosis of diseases, clinical trial designs and drug development.¹⁴ The electrical responses of cancer cell populations are of particular interest, because cancer cells exhibit a collective behavior in terms of cell migration and invasion.^{17–19} However, sensing medium cell populations is extremely challenging because traditional analytical methodologies can be labor-intensive and the compositions of cell monolayers tend to be different. Furthermore, the impedance of cell population is highly sensitive to minute changes in the microenvironment, such as the cell media, pH, temperature and air composition.^{20–23}

Using alternating current (AC) pulses can reduce the noise generated from minute changes in microenvironments of the cell layers, since the impedimetric measurements resulting



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Paper

from an AC pulse from 10 to 100 kHz are largely dominated by the biological cells and the changes within them.^{16,24} This is based on the β -dispersion phenomenon, a theory originally hypothesized by Schwan *et al.* The β -dispersion theory describes how biological cell membranes at low frequencies have a significantly larger impedance compared to cell membranes at high frequencies.^{25,26}

Nanomaterials are promising candidates for detecting cancer cells. The utilization of nanomaterials such as nanosheets, nanotubes and nanodots in cytosensors is becoming increasingly popular, since the quantitative or qualitative detection of cells has wide-ranging applications, such as cancer diagnostics,^{27–30} cell monitoring,³¹ drug development, and other applications.^{32–34} With properties that are significantly different from those of the bulk, the addition of nanomaterials enhances the detection sensitivity of these sensors. For example, leukemia cytosensors using gold nanoparticles have been developed by Zheng *et al.*, which utilize the difference in the expression levels of death receptors on leukemia cells.³⁵ Moreover, Tang *et al.* were able to harness the unique properties of Pt@AgNFs, which were immobilized on a substrate to create circulating-tumor-cell cytosensors.³⁶

Molybdenum disulfide (MoS₂) has emerged as a leading contender for enhancing cancer cell detection due to its unique electrical properties.^{37–39} A two-dimensional (2D) MoS₂ nanosheet comprises a few S–Mo–S monolayers assembled *via* van der Waals (vdW) forces between the layers.³⁷ However, MoS₂ can be utilized in the biomedical field in its early stages due to the limitations imposed by the cytotoxicity of the material and the decreased stability of MoS₂ in aqueous solutions for some applications.⁴⁰ For instance, MoS₂ nanosheets may aggregate in water due to its strong hydrophobic nature and increased vdW attractions between the layers.^{28,41,42}

Herein, we show that by stimulating and altering current flow effects, we can control the electrical conduction of cancer cells via combined 2D-material-polymer-phage frameworks. We then developed an AC-pulse sensor system by using MoS₂/ polyethylene glycol (PEG)/M13 nanoprobes for targeted pancreatic cancer cell (PANC-1) and breast cancer cell (MCF-7) detection. This method achieved a low limit of detection (LOD) of ~15 cells per μ L, which is below an average of \sim 45 cells per µL for existing sensing methods using medium cell population. This allows cancer detection to be performed with a better accuracy for enhancing diagnosis reliability. Additionally, excellent cell viability was achieved in PANC-1 cells at nanoprobe concentrations of up to 50 vol%, which exceeds those utilized in sensing experiments. Furthermore, a nanoprobe concentration of ~ 10 vol%/7.1 nM was attained, below an average probe concentration of ~ 195 nM for current sensing methods with medium cell populations. And, this enables the use of a smaller amount of material for saving diagnosis costs. This proposed methodology for cancer cell detection holds intriguing potential for developing multiplexed lab-on-chip platforms, which can be further applied for clinical purposes such as cancer diagnosis, as well as broad applications in analyte-based sensing.

Methods

Cell lines and cell culture

The PANC-1 cell line was purchased from American Type Culture Collection (ATCC). The PANC-1 cells were cultured in Dulbecco's modified Eagle's medium (DMEM, Nacalai Tasque) supplemented with 10% fetal bovine serum (FBS, Thermo Fisher), 1% L-glutamine (Sigma Aldrich) and 1% penicillinstreptomycin (Fisher Scientific). To elucidate the selectivity of the nanoprobe-type cancer cell system, the MCF-7 and MCF-10A cells were also utilized. The MCF-7 cells were cultured in phenol-red-containing DMEM (Nacalai Tesque), supplemented with 7.5% FBS (Thermo Fisher) and 2 mM of L-glutamine (Sigma Aldrich). The healthy breast epithelial cell line MCF-10A was cultured in DMEM supplemented with Ham's F12 (DMEM/F12, Gibco) containing 10% FBS, 20 ng mL⁻¹ epidermal growth factor (EGF, Gibco), 0.5 μ g mL⁻¹ hydrocortisone (Sigma Aldrich) and 10 μ g mL⁻¹ insulin (Sigma-Aldrich). The cells were incubated at 37 °C in a humidified incubator in an atmosphere of 5% CO₂.

M13 phage propagation

Escherichia coli (E. coli) bacteriophage M13 15669-B1 (M13 phage) was purchased from ATCC. M13 phage was revived according to the phage recovery and propagation protocol (ATCC). 5-Alpha F'Iq competent E. coli (high efficiency E. coli) purchased from New England Biolabs (NEB) was utilized as the host cell for M13 phage propagation. An overnight culture (O.C.) of E. coli was prepared with tetracycline (TET, Sigma Aldrich) and left to incubate on a shaker at 37 °C for 4-6 h at 90 rpm until the mixture showed a cloudy color with an optical absorbance of 0.4 (OD600). A 48 h incubation was then performed with the O.C. and Lennox L broth base (LB Broth, Sigma Aldrich). Similarly, a new culture (N.C.) was prepared and left to incubate at 37 °C for 48 h on a shaker at 90 rpm. First and second precipitations were achieved according to the M13-amplification-protocol (NEB). Concentration of the M13 phage was measured using a µDrop plate (Thermo Fisher Scientific Inc.).

Molecular probe conjugation

Molecular probe conjugation was performed under sterile conditions. MoS_2 (92 mg L⁻¹) in DI water was purchased from 2D Semiconductors Inc. The MoS_2 solution was sonicated prior to molecular probe conjugation. LA-PEG-NHS was purchased from Nanocs Inc. and reconstituted with DI water. The LA-PEG-NHS solution was added to a sterile glass bottle containing MoS_2 in DI water. The mixture was left to incubate on a shaker at 25 °C for 48 h at 120 rpm. The filtered M13 phage was added to the mixture and was then left to incubate on a shaker at 25 °C for 48 h at 120 rpm.

Material characterization

Atomic force microscopy (AFM) was performed using a Bruker Dimension Icon (Bruker Cooperation) system with a scanning area of 1.5 μ m × 1.5 μ m. Probe samples were drop casted on Si

Materials Advances

substrates prior to imaging. Raw AFM data were analyzed with NanoScope analysis software and ImageJ software. Fourier-transform infrared (FTIR) analysis of the probe was performed using Nicolet IS10, US (Thermo Fisher Scientific Inc.) with the wavenumber in the 400–4000 cm⁻¹ range. Transmission electron microscopy (TEM) was carried out using FEI Talos F200 (Thermo Fisher Scientific Inc.). Probe samples were negatively dyed prior to TEM imaging. UV-vis absorbance spectroscopy was performed with a Variskan Lux multimode microplate reader (Thermo Fisher Scientific Inc.).

Cell viability studies

The indium tin oxide (ITO) subsystems were sterilized with 100% ethanol for 20 min prior to exposure to ultra-violet (UV) irradiation for 20 min. The subsystem comprised two left and right ITO electrodes on a glass substrate (Latech) with a cloning cylinder (C3983-50EA, Sigma Aldrich) secured using a silicone adhesive. The size of the gap between the electrodes was chosen to be ~ 0.1 mm, and the thickness of the electrode was fixed at 650 nm. PANC-1 cells were plated in each ITO subsystem at a seeding density of 5×10^3 cells per well and the cells were left to incubate for 24 h. Different concentrations of MoS₂ or MoS₂/ PEG/M13 nanoprobes were added to each subsystem and the cells with the MoS₂ or MoS₂/PEG/M13 nanoprobes were incubated for 24-48 h at 37 °C in a humidified incubator in an atmosphere of 5% CO2. The cytotoxicity of MoS2 was examined using WST-1 assays at 24 h and 48 h after adding MoS₂. Similarly, by utilizing WST-1 assays at 24 h after inserting the nanoprobes, the cytotoxicity of MoS₂/PEG/M13 nanoprobes was evaluated. The absorbance was measured at a wavelength of 450 nm.

Electrical characterization

The ITO subsystems were sterilized with 100% ethanol for 20 min prior to exposure to UV irradiation for 20 min. The PANC-1/MCF-7/MCF-10A cells were plated in each subsystem at different seeding densities (1×10^3 – 7×10^3 cells per well) and the cells were left to incubate for 24 h. 10 vol% MoS₂/PEG/M13 nanoprobes was added to each subsystem. The cells with MoS₂/PEG/M13 nanoprobes were then incubated for 24 h at 37 °C in a humidified incubator in an atmosphere of 5% CO₂. Cell impedance was measured using an arbitrary waveform generator (Tektronix Inc.) and a digital oscilloscope (Tektronix Inc.). Different sensing voltages in the 0.2–0.7 V range were utilized, and the pulse length was fixed at 1 ms.

Results

Methodology of the MoS₂/PEG/M13 sensor framework

The strategy adopted for the highly sensitive detection of cells on the MoS₂/PEG/M13 sensor system is illustrated in Fig. 1. A glass substrate was utilized as the starting material, on which two 650 nm-thick left and right ITO electrodes were deposited for external circuitry connection (Fig. 1(a)). The distance between the electrodes was chosen to be ~100 μ m. The PANC-1/MCF-7 cell layer was then plated. Finally, the MoS₂/ PEG/M13 nanoprobes were added to complete the system structure. Fig. 1(b) shows the variation of normalized system impedance for different probe concentrations. The systems exhibit a decreasing normalized impedance (from 1 to 0.5) with an increase in nanoprobe concentrations (between 0 and 70%). Thus, the electrical conductivity of the sensor system can



Fig. 1 Fabrication of a cancer-cell sensor system integrated with $MoS_2/PEG/M13$ nanoprobes. (a) Schematic illustration of the system structure. (b) Normalized impedance as a function of probe concentration at ~24 h after treatment. Asterisks show the statistical significance of difference in values between the control and test systems (** $p \le 0.01$ and **** $p \le 0.0001$). Data are expressed as the standard error of mean (SEM) where n = 6. (c), (d) Schematic representations of the (c) disulfide binding between the LA terminal of PEG and MoS₂ and (d) amine reaction between pVIII protein of the M13 phage and the NHS terminal of PEG.

Paper

be controlled by varying nanoprobe concentrations. The M13 phage acts as the cell targeting agent, while MoS_2 nanosheets play the role of an impedance contrast agent. PEG comprises lipoic acid (LA) and *N*-hydroxysuccinimide (NHS) ester branches, which can be harnessed to create a conjugated $MoS_2/M13$ system to link MoS_2 nanosheets to the M13 phage.⁴³ The LA-terminal of the PEG linker chemically binds to the MoS_2 nanosheet *via* disulfide binding (Fig. 1(c)).^{44–46} As shown in Fig. 1(d), through amine reactions, the pegylated MoS_2 nanosheets then attach to the M13 phage.⁴⁷ Furthermore, experiments have demonstrated that the pegylation process enhances the stability of MoS_2 nanosheets dispersed in water.^{40,48}

M13 is a cylindrical, ~ 880 nm long and ~ 6.6 nm wide in diameter, nonlytic bacteriophage composed of proteins.⁴⁷ The uptake of M13 phage by cells has been attributed to different pathways of endocytosis.49 Moreover, experiments have revealed that the M13 phage shows preferential binding to the membranes of epithelial cells, and is internalized mainly through clathrin-mediated-endocytosis.50 When left at room temperature, MoS₂ nanosheets undergo spontaneous desulfurization, creating defect sites at the edges of nanosheets.⁵¹ This results in surface electron accumulation, which is substantially larger compared to that in the bulk.⁵¹ The electrical conduction of MoS₂ nanosheets is also stronger than that of bulk materials. Moreover, experiments have demonstrated that surface functionalization of MoS₂ with LA enhances the conduction of MoS₂ nanosheets.⁵² When LA conjugation is utilized, the current flow in MoS₂ nanosheets increases significantly due to enhanced electron mobility. This can facilitate the electrical conductivity of MoS₂/PEG/M13 and improve the sensitivity of nanoprobe sensor systems. Experiments have also revealed that the M13 phage assembles well on cell surfaces.49,53 Moreover, high electrical conductance in cancer cells incubated with MoS₂ is disclosed in recent studies.54,55 Based on these findings and to achieve strong assembly interactions and high electrical conductivity, we chose to utilize the M13 phage and MoS₂ in this work.

Preparation of the MoS₂/PEG/M13 nanoprobe

Liquid-phase exfoliation is an effective method for preparing few-layer MoS_2 nanosheets by breaking weak vdW forces between the layers.^{56,57} We obtained few-layer MoS_2 nanosheets using sonication liquid exfoliation. PEG was then added. The PEG links MoS_2 nanosheets through covalent disulfide bonds between sulphur atoms on LA terminals and exposed sulphur atoms on MoS_2 nanosheets.⁵⁸ Finally, the M13 phage was added to complete the $MoS_2/PEG/M13$ -nanoprobe-structure. The PEGylated MoS_2 binds to the M13 phage in an NHS ester-amine reaction to form an amide bond between PEG and M13 phage.⁴³

The chemical composition and morphology of the prepared samples were systematically investigated. Fig. 2(a) shows the transmission electron microscopy (TEM) image of the $MoS_2/PEG/M13$ nanoprobe, in which thin 2D nanosheets and 1D wire-like structures were observed. To investigate the crystal

structure and quality of the nanoprobe, the Fourier-transform infrared (FTIR) spectra of the MoS₂/PEG/M13 nanoprobe sample were obtained (Fig. 2(b)). A broad peak at \sim 3400 cm⁻¹ was observed in the FTIR spectra of the MoS₂/PEG/M13 nanoprobe sample. This corresponds to an amide bond stretching (N-H) between PEG and M13 phage. Another peak at ~ 1640 cm⁻¹ corresponds to a carbonyl stretch (C=O) associated with amide bonds. Although amide bonds are present in all biological molecules, including the M13 phage, these bonds are vital in determining the success of probe conjugation. Additionally, the lack of a peak in the 2500–2600 cm^{-1} range, corresponding to a thiol stretch (S-H), indicates that all LA terminals on the PEG linker are covalently bound to the available MoS₂ nanosheets. Additional peaks observed at 607, 875, 1112, 1427 and 1627 cm⁻¹ are characteristic of MoS_2 , 45,59,60 while the peaks at 2800-3000 cm⁻¹ correspond to the presence of PEG.^{61,62} These results indicate that both MoS₂ and PEG are present in the nanoprobes. The atomic force microscopy (AFM) topology of the nanosheet of the MoS₂/PEG/M13 nanoprobe is shown in Fig. 2(c). The thickness of the nanosheet was determined by cross-sectional analysis (Fig. 2(c), line A-B). The measured height of ~ 7 nm shown in Fig. 2(d) corresponds to MoS₂ nanosheets with ~ 10 triple layers.⁶³

Ultraviolet-visible (UV-vis) absorbance spectroscopy was further performed to investigate the presence of the M13 phage and MoS₂ in the conjugated MoS₂/PEG/M13 nanoprobe system (Fig. S1, ESI[†]). Fig. S1a (ESI[†]) shows that both MoS₂ and MoS₂/ PEG/M13 samples exhibit similar spectral features. However, the absorbance intensity of the MoS₂/PEG/M13 sample is smaller compared to that of the MoS₂ sample. The absorbance intensity may correlate to the concentration of MoS₂. As MoS₂ is hydrophobic, the conjugation process can result in an increased dispersion of MoS₂, resulting in a decrease in the absorbance intensity of the MoS₂/PEG/M13 sample. A signature peak at ~ 250 nm was observed for both M13 and MoS₂/PEG/ M13 samples, similar to those reported in the literature.^{44,64,65} This indicates that the conjugation process does not alter spectral characteristics of the M13 phage.

The viability of PANC-1 cells with pure MoS₂ was investigated at different material concentrations using WST-1 cell viability assay after 24 h and 48 h incubation periods (Fig. 2(e)). The cells with 25–100 μ M MoS₂ disclose similar cell viability to that of the control (cells only/0 µM MoS₂), indicating that MoS_2 has negligible cytotoxic effects. Additionally, we investigated the viability of PANC-1 cells with MoS₂/PEG/M13 nanoprobes 24 h after incubation (Fig. 2(f)). The cells with 10-50 vol% MoS₂/PEG/M13 nanoprobes exhibit a similar cell viability of $\sim 100\%$ compared to that of the control (0 vol%) MoS₂/PEG/M13 nanoprobes), whereas a decreased cell viability of $\sim 60\%$ compared to that of the control was shown by cells with $\sim 70 \text{ vol}\% \text{ MoS}_2/\text{PEG}/\text{M13}$ nanoprobes. For bare MoS_2 , small/large material concentrations can maintain high cell viability, as shown in experiments. However, for the MoS₂/ PEG/M13 nanoprobe, the charge level of the nanoprobe can reach a value that induces cell death when nanoprobes are incubated with cells.32 However, the concentration of the



Fig. 2 Characterization of MoS₂/PEG/M13 nanoprobes. (a) Transmission electron microscopy (TEM) image of the MoS₂/PEG/M13 nanoprobe, showing thin 2D nanosheets and 1D wire-like structures. (b) Fourier-transform infrared (FTIR) spectra of MoS₂/PEG/M13-nanoprobe samples. The FTIR spectra were normalized to the maximum absorbance value. (c) Atomic force microscopy (AFM) image of a nanosheet of the MoS₂/PEG/M13 nanoprobe. (d) Height profile of the MoS₂ nanosheet along the line A–B in (c). (e), (f) Viability of PANC-1 cells with (e) pure MoS₂ and (f) MoS₂/PEG/M13 nanoprobes determined *via* WST-1 assay. The absorbance was measured at $\lambda = 450$ nm. Cells were grown on ITO systems ~24 h prior to the addition of MoS₂ or MoS₂/PEG/M13 nanoprobes. Asterisks indicate the statistical significance of difference in values between the control and test systems (*** $p \le 0.001$, **** $p \le 0.0001$ and non-significant (ns)). Data are expressed as the standard error of mean (SEM) where n = 6.

nanoprobe is not small enough to prevent cell death at high nanoprobe concentrations, leading to low cell viability.

Cell sensing performance of MoS₂/PEG/M13 nanoprobes

We measured and calculated the normalized impedance of the PANC-1/MCF-7 cell systems with $MoS_2/PEG/M13$ nanoprobes for different cell populations/sensing voltages. The impedance of the system was normalized to that of the control (cells only). The PANC-1 cell system with 10 vol% $MoS_2/PEG/M13$ nanoprobes showed ~16% decrease in impedance compared to that of the control, whereas ~50% smaller impedance than that of the control was observed for the PANC-1 system with 70 vol% $MoS_2/PEG/M13$ nanoprobes (Fig. 1(b)). To increase the conductance, and at the same time, maintain excellent cell viability, and also minimal nanoprobe concentrations, we chose to

utilize the PANC-1 cell system with 10 vol% MoS₂/PEG/M13 nanoprobes. Utilizing the PANC-1 cell system with 10 vol% MoS₂/PEG/M13 nanoprobes, we observed a decrease in the normalized impedance (the normalized impedance is defined as the ratio of the impedance of the cell system with MoS₂/PEG/M13 nanoprobes to the impedance of a pure cell system) for increasing cell populations (Fig. 3(a)). When 3×10^3 and 5×10^3 cells were utilized, we observed a corresponding ~14% and ~25% decrease in the normalized impedance. The largest percentage decrease in normalized impedance was observed for 7×10^3 cells (above 30%).

To investigate the LOD of the PANC-1 cell system with 10 vol% $MoS_2/PEG/M13$ nanoprobes, we performed a linear regression analysis based on the slope of the linear plot of the normalized impedance as a function of the cell population



Fig. 3 Sensing cancer cells using a MoS₂-PEG-M13-nanoprobe-based AC-pulse sensor system. (a) Cell population dependent normalized impedance values for PANC-1 cells. (b) Sensing voltage dependence of normalized impedance for PANC-1 cells. The cell population is fixed at 5×10^3 cells. (c), (d) Normalized impedance variations for different cell populations for (c) MCF-7 and (d) MCF-10A cells. Asterisks indicate the statistical significance of difference in values between the control and test systems (* $p \le 0.05$, ** $p \le 0.01$, **** $p \le 0.0001$ and non-significant (ns)). Data are expressed as the standard error of mean (SEM) where n = 6.

(gradient of the dotted line in Fig. 3a) (s), and the standard deviation (σ) of the normalized impedance:

$$\text{LOD} = 3.3 \times \left(\frac{\sigma}{s}\right) \tag{1}$$

The calculated LOD was found to be ~15 cells per μ L $\left(\text{LOD} = 3.3 \times \frac{2.0 \times 10^{-4}}{4.5 \times 10^{-5}} = 15.2\right)$, below the baseline of ~50 cells per μ L for state-of-the-art sensing methods⁶⁶⁻⁷⁰ with medium cell populations (Fig. S2, ESI†). Moreover, the concentration of the MoS₂/PEG/M13 nanoprobe utilized for the LOD analysis was ~10 vol%/7.1 nM, which is lower than the average probe concentration of ~195 nM for existing sensing methods⁷¹⁻⁷³ with medium cell populations (Fig. S3, ESI†). Additionally, the normalized impedance of the PANC-1 cell system with 10 vol% MoS₂/PEG/M13 nanoprobes decreases (from 0.7 to 0.6) with an increase in sensing voltage (0.2–0.7 V) (Fig. 3(b)). These results indicate that the electrical conductance of nanoprobe-based cell systems can be tuned for sensitive cancer cell detection.

In cancer cell sensing studies, the ability to distinguish cancer cells from normal cells is an important aspect. Experiments have demonstrated that nanomaterial-based platforms disclose the ability to distinguish cancer cells from normal cells for different types of organs or between different cell types. The MoS₂-based framework with cervical and lung cancer blood samples reveals a different relative impedance as compared

with that of the framework with healthy cervical and lung blood samples.⁷⁴ A lower impedance is exhibited in a gold nanoparticle-based system with HeLa cells compared to the system with human embryo skin fibroblast cells (HESF cells).75 The graphene-based platform with HeLa cells also shows a smaller impedance than that of the platform with NIH3T3 cells.⁷⁶ Furthermore, we compared the normalized impedance of the MCF-7 cell system with 10 vol% MoS₂/PEG/M13 nanoprobes with that of the MCF-10A cell system with 10 vol% MoS₂/PEG/ M13 nanoprobes, albeit utilizing different cell lines (Fig. 3(c) and (d)). When the MCF-7 cell system with MoS₂/PEG/M13 nanoprobes for moderate cell populations (*i.e.*, 3×10^3 and 5×10^3 cells) was examined, the normalized impedance (0.7– 0.8) was found to be smaller relative to that of the MCF-10A cell system with $MoS_2/PEG/M13$ nanoprobes (~1.0), indicating that MCF-7 cells are more sensitive/selective to the MoS₂/PEG/M13 nanoprobe than MCF-10A cells in terms of sensing ability.

Discussion

Electrical-based sensing is a popular approach for detecting cancer cells, since cell types have unique electrical signatures that can be harnessed for disease diagnosis.^{16,77–80} In disease diagnosis, cell counts are clinically relevant, where their utilization has been adopted as a predictor of patient prognosis, such as lymphocyte counts in patients with lung cancer,⁸¹ monocyte

8

counts in malignant-pleural-mesothelioma,⁸² and white-bloodcell counts for other types of cancers.^{83–85} In this work, the systems with MoS₂/PEG/M13 nanoprobes exhibit a low LOD of ~15 cells per μ L, below the reference of ~50 cells per μ L for current sensing methods with medium cell populations. This was achieved by enhancing current flow through the cells, and at the same time, maintaining excellent cell viability.

Moreover, achieving rapid responses in sensor systems is critical for providing key information that could aid in patient management. Traditional sensors may exhibit a detection time in the 1–24 h range.^{68,86} In this work, the impedance could be obtained using millisecond pulses, enabling real-time monitoring of cancer cells. Additionally, the setup is facile and most of the components in the system are reusable, which can save diagnosis-related costs.

In the future, work can be performed to utilize the multiplexed potential of the M13 phage to match the complexity of cancer as a disease. The high density of available functionalization sites at the pIII and pVIII protein sites has not been fully utilized to broaden the capabilities of MoS₂/PEG/M13 nanoprobes for improved targeting and detection.47 Other nanomaterials/phages could also be utilized to optimize cancer cell sensor performance. Experiments have shown that biological systems with other nanomaterials, viz., metal nanoparticles and nanowires, reveal a low LOD.^{87,88} Additionally, biological platforms with other phages, e.g., the Listeria phage and T7 bacteriophage, which disclose a low LOD, have been demonstrated in recent studies.^{89,90} These experiments elucidate that the nanomaterial/phage-based interaction facilitated an increase in sensitivity. Application of the MoS₂/PEG/M13 nanoprobes can be further explored in animal studies and clinical trials, with different types of cancers. Oncologists may rely on traditional morphological or histological characteristics of tumor tissues for disease diagnosis, which cannot be determined affirmatively with a single type of analysis. Tumor development is an ongoing biological process,91 due to which the validity of these analyses is short-lived.⁹² Multi-parametric testing using the MoS₂/PEG/M13 nanoprobes open a path to enhance the sensitivity and speed of diagnosis for improving patient well-being.

In recent years, there has been a gradual shift in understanding the developmental behavior of cancer. A growing number of studies have suggested that the predominant mode of cancer invasion is through collective cell migration,¹⁷ where cells move as clusters, strands or sheets. Collective cell migration has become a hallmark of cancer metastasis in many types of tumor,¹⁹ implying that traditional single-cell analyses may no longer be sufficient for diagnosing and understanding the cancer-cell behavior accurately.93 Furthermore, the bioelectrical properties of cancer cells have been demonstrated to affect not just its own behavior, but also the behavior of neighboring cells.⁹⁴ With the growing association between bioelectrical properties and collective cell behaviors demonstrated in cancer cells, impedimetric cell-population-based methods of detection can provide valuable information in understanding cancer cell development.

Conclusion

In this work, we demonstrated an ultra-sensitive and low-probeconcentration MoS₂/PEG/M13-nanoprobe-based AC-pulse sensor system for detecting PANC-1 and MCF-7 cells. This is achieved through combined 2D-material-polymer-phage frameworks that alter the electrical conduction of cancer cells. The MoS₂/PEG/M13-nanoprobe-based PANC-1 cell system has an LOD of ~15 cells per μ L, below the reference value of ~ 50 cells per µL for current sensing methods using medium cell population. In addition, excellent cell viability was achieved in PANC-1 cells with a MoS₂/PEG/M13 nanoprobe concentration up to 50 vol%, which exceeds the concentrations utilized in detection experiments. Furthermore, a nanoprobe concentration of ~10 vol%/7.1 nM was achieved, below an average of ~ 195 nM used for existing sensing methods with medium cell populations. As a result, the proposed system represents the first methodology reported using MoS₂/PEG/ M13 nanoprobes for clinically relevant cancer cell detection and constitutes an excellent opportunity for the development of lab-on-chip platforms.

Data availability

The authors declare that the data supporting the finding of this study are available within the article and the ESI.† Other data are available from the corresponding authors upon reasonable request.

Conflicts of interest

The authors declare no competing interests.

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